IN THE UNITED STATES DISTRICT COURT FOR THE WESTERN DISTRICT OF PENNSYLVANIA

IN RE: PHILIPS RECALLED CPAP,

BI-LEVEL PAP, AND MECHANICAL

VENTILATOR PRODUCTS

This Document Relates to:

LITIGATION

Jamison Drake

MDL No. 3014

SHORT FORM COMPLAINT FOR

Master Docket: Misc. No. 21-mc-1230-JFC

PERSONAL INJURIES, DAMAGES,

AND DEMAND FOR JURY TRIAL

2:23-CV-318

Judge Conti

Plaintiff(s) incorporate(s) by reference the Amended Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial filed in In re Philips Recalled CPAP, Bi-Level PAP, and Mechanical Ventilator Products Litigation, MDL No. 3014, Master Docket Misc. No. 21-mc-1230 (the "Master Long Form Complaint"). This Short Form Complaint adopts the allegations, claims, and requested relief as set forth in the Master Long Form Complaint. As necessary herein, Plaintiff(s) may include: (a) additional claims and allegations against Defendants; and/or (b) additional claims and allegations against other Defendants not listed in the Master Long Form Complaint.

Plaintiff(s) further allege(s) as follows:

DEFENDANTS I.

- Plaintiff(s) name(s) the following Defendants in this action: 1.
 - Koninklijke Philips N.V.
 - Philips North America LLC.
 - Philips RS North America LLC.

		Philips Holding USA Inc.			
		Philips RS North America Holding Corporation.			
		Polymer Technologies, Inc.			
		Polymer Molded Products LLC.			
II.	PLAINTIFF(S)				
	2.	Name of Plaintiff(s): Jamison Drake			
	3.	Name of spouse of Plaintiff (if loss of consortium claim is being made): N/A			
	4.	Name and capacity (<i>i.e.</i> , executor, administrator, guardian, conservator, etc.) of other Plaintiff, if any: N/A			
	5.	State(s) of residence of Plaintiff(s) (if the Recalled Device user is deceased residence at the time of death): Texas			
III.	DES :	DESIGNATED FORUM 6. Identify the forum (United States District Court and Division) in which the Plainting			
		would have filed in the absence of direct filing: Texas Northern District Court – Fort Worth, TX			

IV. USE OF A RECALLED DEVICE

7. Plaintiff used the following Recalled Device(s):

E30 (Emergency Use Authorization)	Dorma 500		
DreamStation ASV	REMstar SE Auto		
☐ DreamStation ST, AVAPS	Trilogy 100		
SystemOne ASV4	Trilogy 200		
C-Series ASV	Garbin Plus, Aeris, LifeVent		
C-Series S/T and AVAPS	A-Series BiPAP Hybrid A30 (not marketed		
OmniLab Advanced +	in U.S.)		
SystemOne (Q-Series)	A-Series BiPAP V30 Auto		
✗ DreamStation	A-Series BiPAP A40		
DreamStation Go	A-Series BiPAP A30		
Dorma 400	Other Philips Respironics Device; if other,		
	identify the model:		
V. INJURIES			
· ·	g physical injuries as a result of using a Recalled dant symptoms and consequences associated		
COPD (new or worsening)			
X Asthma (new or worsening)			
Pulmonary Fibrosis			
Other Pulmonary Damage/Inflammatory Response			
Cancer	(specify cancer)		
Kidney Damage			
Liver Damage			

I	Heart Damage
I	Death
	Other (specify) Bronchitis

VI. CAUSES OF ACTION/DAMAGES

9. As to Koninklijke Philips N.V., Plaintiff(s) adopt(s) the following claims asserted in the Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial, and the allegations and prayer for relief with regard thereto, as set forth therein:

Count I: Negligence

Count II: Strict Liability: Design Defect

Count III: Negligent Design

Count IV: Strict Liability: Failure to Warn

Count V: Negligent Failure to Warn

Count VI: Negligent Recall

Count VII: Battery

Count VIII: Strict Liability: Manufacturing Defect

Count IX: Negligent Manufacturing

Count X: Breach of Express Warranty

Count XI: Breach of the Implied Warranty of Merchantability

Count XII: Breach of the Implied Warranty of Usability

Count XIII: Fraud

Count XIV: Negligent Misrepresentation

X Count XV: Negligence Per Se Count XVI: Consumer Fraud and/or Unfair and Deceptive Practices Under State Law Count XVII: Unjust Enrichment Count XVIII: Loss of Consortium Count XIX: Survivorship and Wrongful Death Count XX: Medical Monitoring Count XXI: **Punitive Damages** Count XXII: Other [specify below] As to Philips North America LLC, Plaintiff(s) adopt(s) the following claims asserted in the Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial, and the allegations and prayer for relief with regard thereto, as set forth therein: Count I: Negligence Count II: Strict Liability: Design Defect **X** Count III: Negligent Design **X** Count IV: Strict Liability: Failure to Warn **X** Count V: Negligent Failure to Warn Count VI: Negligent Recall **X** Count VII: Battery Count VIII: Strict Liability: Manufacturing Defect Count IX: Negligent Manufacturing

10.

Count X: Breach of Express Warranty **X** Count XI: Breach of the Implied Warranty of Merchantability Count XII: Breach of the Implied Warranty of Usability Count XIII: Fraud Count XIV: Negligent Misrepresentation Count XV: Negligence Per Se Count XVI: Consumer Fraud and/or Unfair and Deceptive Practices Under State Law Count XVII: Unjust Enrichment Count XVIII: Loss of Consortium Count XIX: Survivorship and Wrongful Death Count XX: Medical Monitoring **X** Count XXI: **Punitive Damages** Count XXII: Other [specify below] As to Philips RS North America LLC, Plaintiff(s) adopt(s) the following claims

11. asserted in the Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial, and the allegations and prayer for relief with regard thereto, as set forth therein:

Count I: Negligence

Count II: Strict Liability: Design Defect

X Count III: Negligent Design

Count IV: Strict Liability: Failure to Warn

X Count V:	Negligent Failure to Warn
X Count VI:	Negligent Recall
Count VII:	Battery
Count VIII:	Strict Liability: Manufacturing Defect
X Count IX:	Negligent Manufacturing
Count X:	Breach of Express Warranty
X Count XI:	Breach of the Implied Warranty of Merchantability
X Count XII:	Breach of the Implied Warranty of Usability
Count XIII:	Fraud
Count XIV:	Negligent Misrepresentation
Count XV:	Negligence Per Se
X Count XVI:	Consumer Fraud and/or Unfair and Deceptive Practices Under State Law
X Count XVII:	Unjust Enrichment
Count XVIII:	Loss of Consortium
Count XIX:	Survivorship and Wrongful Death
Count XX:	Medical Monitoring
Count XXI:	Punitive Damages
Count XXII:	Other [specify below]

12.

12.	in the Master Long	ng USA Inc., Plaintiff(s) adopt(s) the following claims asserted Form Complaint for Personal Injuries, Damages and Demand he allegations and prayer for relief with regard thereto, as set
	Count I:	Negligence
	Count II:	Strict Liability: Design Defect
	X Count III:	Negligent Design
	X Count IV:	Strict Liability: Failure to Warn
	X Count V:	Negligent Failure to Warn
	X Count VI:	Negligent Recall
	X Count VII:	Battery
	X Count VIII:	Strict Liability: Manufacturing Defect
	X Count IX:	Negligent Manufacturing
	Count X:	Breach of Express Warranty
	X Count XI:	Breach of the Implied Warranty of Merchantability
	Count XII:	Breach of the Implied Warranty of Usability
	Count XIII:	Fraud
	X Count XIV:	Negligent Misrepresentation
	Count XV:	Negligence Per Se
	Count XVI:	Consumer Fraud and/or Unfair and Deceptive Practices Under State Law
	Count XVII:	Unjust Enrichment
	Count XVIII:	Loss of Consortium
	Count XIX:	Survivorship and Wrongful Death
	X Count XX:	Medical Monitoring

Count XXI: **Punitive Damages** Count XXII: Other [specify below] As to Philips RS North America Holding Corporation, Plaintiff(s) adopt(s) the following claims asserted in the Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial, and the allegations and prayer for relief with regard thereto, as set forth therein: Count I: Negligence **X** Count II: Strict Liability: Design Defect **X** Count III: Negligent Design **X** Count IV: Strict Liability: Failure to Warn Count V: Negligent Failure to Warn Count VI: Negligent Recall Count VII: **Battery**

Count X: Breach of Express Warranty

Count XI: Breach of the Implied Warranty of Merchantability

Strict Liability: Manufacturing Defect

Count XII: Breach of the Implied Warranty of Usability

Count XIII: Fraud

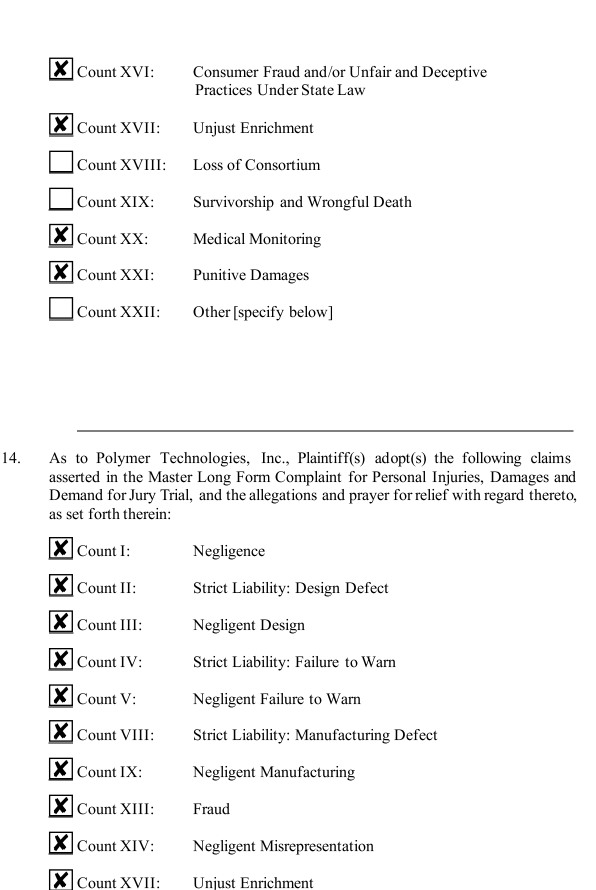
X Count VIII:

X Count IX:

13.

Count XIV: Negligent Misrepresentation

Count XV: Negligence Per Se



	Count XVIII:	Loss of Consortium
	Count XIX:	Survivorship and Wrongful Death
	Count XX:	Medical Monitoring
	Count XXI:	Punitive Damages
	Count XXII:	Other [specify below]
15.	asserted in the Mast	ded Products LLC, Plaintiff(s) adopt(s) the following claims are Long Form Complaint for Personal Injuries, Damages and al, and the allegations and prayer for relief with regard thereto
	X Count I:	Negligence
	X Count II:	Strict Liability: Design Defect
	Count III:	Negligent Design
	X Count IV:	Strict Liability: Failure to Warn
	X Count V:	Negligent Failure to Warn
	Count VIII:	Strict Liability: Manufacturing Defect
	X Count IX:	Negligent Manufacturing
	Count XIII:	Fraud
	Count XIV:	Negligent Misrepresentation
	X Count XVII:	Unjust Enrichment
	Count XVIII:	Loss of Consortium
	Count XIX:	Survivorship and Wrongful Death
	Count XX:	Medical Monitoring

18. Plaintiff(s) assert(s) the following additional claims and factual allegations against other Defendants named in Paragraph 16 above:

WHEREFORE, Plaintiff(s) pray(s) for relief and judgment against Defendants and all such further relief that this Court deems equitable and just as set forth in the Master Long Form Complaint for Personal Injuries, Damages and Demand for Jury Trial and any additional relief to which Plaintiff(s) may be entitled.

Date: <u>Feb</u> 27 2023

/s/ Shanon J. Carson

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